BIOLOGICAL CONTROL

Predation as a Mortality Factor in Populations of the Spittlebug, *Deois flavopicta* Stål (Homoptera: Cercopidae)

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RESUMO - A cigarrinha-das-pastagens *Deois flavopicta* Stål é a principal praga de pastagens cultivadas nos Cerrados do Brasil Central. O inseto apresenta distribuição sazonal com três gerações anuais discretas durante a estação chuvosa (setembro-abril) e população sincronizada de ovos diapáusicos durante a estação seca (maio-agosto). Experimentos em pastagens cultivadas mostraram que predadores não-específicos foram capazes de alterar significativamente as taxas de mortalidade de ovos diapáusicos e ninhas de *D. flavopicta*. Predação foi um fator de mortalidade independente da densidade, reduzindo os ovos diapáusicos em aproximadamente 60% e ninhas de 20% a 47%. A taxa de mortalidade das ninhas resultante da predação não variou significativamente entre a primeira e a segunda geração anual. No entanto, a mortalidade de ninhas na terceira geração foi significativamente mais baixa, fato relacionado ao maior grau de agregação das ninhas em refúgios espaciais. Observações diretas de interações predador-presa nas pastagens sugerem que, potencialmente, as formigas representam o grupo que mais contribui para a mortalidade observada, embora esse impacto deva ainda ser verificado experimentalmente. Taxas de mortalidade mais altas em ovos e ninhas de *D. flavopicta* expostos aos predadores indicam que a predação pode ser um fator importante na determinação da população de adultos. Desta forma, práticas de manejo das pastagens que perturbam a comunidade de predadores podem resultar em densidades mais altas das cigarrinhas.

PALAVRAS-CHAVE: Insecta, comunidade de insetos, dinâmica populacional, curva de sobrevivência

ABSTRACT - The spittlebug, *Deois flavopicta* Stål, is the main pest in cultivated pastures of the “Cerrados” (savanna) in the central region of Brazil. The insect has three discrete generations during the rainy season (September-April) and a synchronized population of diapausuring eggs during the dry season (May-August). Experiments in cultivated pastures showed that nonspecific predators were able to affect significantly the mortality rates of diapausuring eggs and nymphs of *D. flavopicta*. Predation was a density independent mortality factor that reduced diapausuring eggs by approximately 60% and nymphs by 20% to 47%. The nymph mortality rate due to predation did not differ during the first and the second generations of the year. For the third generation however, the mortality rate was lower, related to a greater degree of nymphal aggregation in spatial refuges. Our direct observations indicate that among the types of predators in pastures, ants potentially make the greatest contribution to mortality, although their impact needs experimental verification. High rates of mortality of eggs and nymphs of *D. flavopicta* exposed to predators indicate that predation can be an important factor determining the size of adult populations. Therefore, management practices that disturb the predator community may increase the population densities of the spittlebug.

KEY WORDS: Insecta, insect community, population dynamics, survival curve

Since 1960, several grass species of the genus *Brachiaria* (Poaceae) have been introduced from Africa and planted in extensive areas, from the Central to Northern regions of Brazil (Seiffert 1980). The colonization of new pastures with few natural enemies, along with a phenology that is synchronized with the seasonal change in physical
factors and the life cycle of the host plant, has favored frequent spittlebug (Homoptera: Cercopidae) outbreaks in several regions of Brazil (Cosenza & Naves 1980). Spittlebug population levels are controlled by weather variations, intrinsic regulators of the population and by the growth pattern of the host plant (Menezes et al. 1983). While resistant to drought, introduced Brachiaria species did not possess resistance to spittlebugs (Niklakhe 1987, Cosenza et al. 1989).

Among spittlebugs, Deois flavopicta (Stal) is the main species in pasture areas of Central Brazil in the “Cerrados” region. Each year this species has three discrete generations, during the rainy season, and it aestivates during the dry season as diapausing eggs (Fontes et al. 1995). Low temperatures during the early part of the diapause period accelerate diapause development, whereas contact with liquid water determines the timing of post-diapause hatching (Pires et al. 2000). With the start of the rainy season, synchronized emergence of first instars occurs after the natural termination of diapause during the dry season (Sujii et al. 1995).

Various species of insects, spiders, nematodes, and birds are cited as natural enemies of spittlebug (Villacorta 1980; Barbosa et al. 1984; Carneiro 1988; Hewitt & Nilakhe 1986; Bueno 1987; Marques 1988; Pires et al. 1993). Among them, parasitoids of D. flavopicta are few, but include the egg parasitoid Anagrus sp. that attacks ca. 2% of the eggs (Pires et al. 1993). Entomopathogens include Metarhizium anisopliae (Metsch.) Sorokin, Beauveria bassiana (Bals.) Vuillemin, and Paecilomyces lilacinus (Thom.) Samson (Faria & Tiggano 1996), but our field experience and knowledge of this insect indicate that their impact on spittlebug populations is minimal. Predators such as ants (Hymenoptera: Formicidae) (Hewitt & Nilakhe 1986) and Salpingogaster nigra Schiner (Diptera: Syrphidae) (Marques 1988) are mentioned, for some regions, as the main possible causes of mortality that are able to significantly alter the density of spittlebug populations. However, the impact of predators on the regulation of D. flavopicta population has not been experimentally investigated.

Outbreaks of D. flavopicta, common in Central Brazil, cause severe damage to pastures and cattle production. Population-regulating factors are being explored in search for environmentally safe methods for managing this pest. In this paper, we investigated the role of predators in the regulation of D. flavopicta populations. This knowledge will support new proposals for management of pastures to favor natural biological control and reduce the occurrence of economically damaging populations of spittlebugs.

Material and Methods

Experimental Plots. This study was conducted in three experimental plots located at Embrapa/Cenargen, Embrapa/Cerrados, and Granja do Torto, in Brasilia, DF, Brazil. At Embrapa/Cenargen, the experiments were run in a 0.8 ha of pasture cultivated with Brachiaria decumbens Stapf (Poaceae). At Embrapa/Cerrados and Granja do Torto the experiments were conducted in pastures of ca. 3 ha, cultivated with Brachiaria ruziensis Stapf.

Predation of Diapausing Eggs During the Dry Season. Beginning June 1996, groups of diapausing eggs were placed in 15 randomly selected sites within the pasture. To mark the position of the eggs, allow minimal alteration of soil permeability characteristics, and allow access to natural enemies, a 10 cm x 10 cm piece of organdy screen was placed over the soil and groups of diapausing eggs were then placed over these screens. Both screens and eggs were then covered with approximately 1 cm of soil. To test for effects of density on egg removal rates by predators, five groups of 100 eggs and 10 groups of 50 eggs were used. Screens were gathered three months later and the eggs that remained on the screens were collected by filtering, flotation in a saturated NaCl solution and observation under a stereomicroscope (see Sujii 1994). The proportions of missing eggs from the two different density conditions were compared with a “t” test (Wilkinson 1990).

Immature Mortality. Nymphal mortality in pastures was evaluated during the second generation of each year from 1994 to 1997 and during the three generations of 1995/96 (October to May) using a Modified Mark Release and Recapture (MMRR) method. This method consisted of finding, marking position, and releasing all nymphs in 10 randomly selected 0.5 m x 0.5 m plots. Nymphs were marked and followed until adult emergence. A small numbered wooden stake indicated the position of the nymphs in each plot. The developmental stage of nymphs was determined by morphological characteristics, and data related with 3rd and 4th instars were joined because of the difficulty to separate them (Menezes et al. 1983, Sujii 1994). The position and development of the nymphs were recorded every one to three days.

During 1995/96 (three generations) and 1997 (second generation), nymphal cohorts protected against predators were followed through the MMRR method, as previously described. Nymphal development was measured inside 10 randomly located plots protected by a cage measuring 0.5 m x 0.5 m and 0.7 m high. The cages were wooden framed and covered with organdy screen panels to exclude predators. This set of cohorts was used as control for estimating predation in unprotected individuals.

Survival curves from first instar to teneral adults were calculated and compared graphically for each generation. Comparisons of mortality rates among different generations and years were done using Analysis of Variance followed by Tukey’s Test (Wilkinson 1990). The effect of density on mortality rates in protected and unprotected quadrats was evaluated through Pearson’s Correlation Coefficient (Wilkinson 1990). The pattern of nymphal aggregation was calculated through the Morisita’s index of dispersion (Krebs 1989). This index was calculated using the density of nymphs in each unprotected quadrat for the set of quadrats in each generation. This index was correlated with the correspondent mortality to test if predation levels could be related to spatial distribution of nymphs.

Direct Observation and Collection of Predators. To be efficient in performing direct observations of predation, it was necessary to establish the diurnal pattern (day vs. night)
of predator activity. To do so, we determined when nymphs disappeared by following a group of D. flavopicta (all instars) using the MMRR method. Observations were made twice a day at 12h intervals, early in the morning (7:00 am) and in the evening (7:00 pm), for 10 days. Rates of disappearance of nymphs, inferred as predator attack, during each period (night and day) were compared with the chi-square test.

Direct observation of predators was made for ca. 120 min. per sampling along eight transects of 0.5 m x 20 m randomly marked in a pasture of B. ruziziensis at Granja do Torto. During observations, prey and predators seen inside transects were recorded. Predators were followed for five minutes. When an attack on a spittlebug occurred, both prey and predator were counted and collected. Unknown insects and predators that were already attacking prey were also collected for identification. Insect samples were sent to Universidade Estadual de Campinas, Comissão Executiva de Pesquisa da Lavoura Cacaueira, and Universidade de Brasilia for identification. Data were plotted in order to describe the insect community present in the pastures.

Results

Predator Effect on Survival of Diapausing Eggs. During the sampling collection at the end of dry season, no disturbance or water run off was observed in the soil over the diapausing eggs. This observation suggests that missing eggs were mainly caused by action of predators. The percentages of diapausing eggs missing from groups of 50 and 100 eggs per sample were respectively 60.0 ± 26.15% and 62.8 ± 19.61% (mean ± standard deviation), and did not statistically differ (t-test, $t = 0.21$; d.f. = 14; $P = 0.84$). However, the average numbers of missing eggs from plots with 100 eggs (62.8 ± 19.61%) was higher than from plots with 50 eggs (30.0 ± 12.40%), revealing that the number of attacks per plot increased with prey density.

Nymph Survival. For the different generations of 1996, survival curves of uncaged nymphs, mainly first and second generations, had similar general shapes, with mortality occurring early. In contrast, nymphs protected with anti-predator cages had relatively constant rates of mortality and higher overall survivorship rates (Fig. 1). In 1995/96, the good distribution and high levels of precipitation during the period of October-May appear to have favored nymphal survival and produced high spittlebug densities. The average densities of nymphs during the three population peaks were 40, 60 and 20 individuals/m², reaching numbers up to 200 individuals/m² in some plots during the 2nd population peak. Survival curves of the 2nd generation from different years (1994-97) showed similar shapes and rates (Fig. 2).

Nymphs survival rates were higher in the quadrats protected by cages in comparison to the quadrats without cages (Table 1). A total of 1,562 nymphs was observed, 888 in cages and 674 in quadrats without cages, during 1995/96. The densities varied between 3 to 108 nymphs/quadrat (equivalent to 12 to 432 individuals/m²). Of these, 928 individuals reached the adult stage (664 from cages and 264 from quadrats without cages). Average survival rates in the protected quadrats did not differ during 1995/96 and 1997 generations, and were higher than those in the quadrats without cages, in the first and second generations (See ANOVA, Table 1).

For 1995/1996, mortality rates remained constant across the instars during the three annual generations for uncaged populations (Fig. 3). In the caged treatments, no significant correlation between density and proportion of nymphal mortality was found within cages ($r = 0.305, n = 30, P = 0.108$) (Fig. 4b). A significant correlation was observed between nymphal densities and mortality rate, for uncaged treatments (Pearson’s coefficient, $r = 0.525 n = 30, P = 0.003$) (Fig. 4a). However, there was no significant correlation between nymphal density and mortality rates within non-caged treatments when each generation is analyzed independently (Table 2). A negative
Figure 3. Mortality of uncaged spittlebug, *D. flavopicta*, in the Distrito Federal, during 1995/96. No significant differences among instars of the same generation were observed (Kruskal Wallis, test *P* < 0.05).

Table 1. Mortality (± s. dev.) of immature spittlebugs, *D. flavopicta* observed in field quadrats during the rainy season of 1995/96 and 1996/97 in pastures of *B. ruzicrensis* located in Brasilia, DF, Central Brazil.

<table>
<thead>
<tr>
<th>Generation/year</th>
<th>Caged(n)</th>
<th>Uncaged(n)</th>
<th>t Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>1&lt;sup&gt;st&lt;/sup&gt;/1996</td>
<td>13.4 ± 15.55(10) a</td>
<td>57.1 ± 21.89(10) ab</td>
<td>t = 5.147 <em>P</em> &lt; 0.001</td>
</tr>
<tr>
<td>2&lt;sup&gt;nd&lt;/sup&gt;/1996</td>
<td>30.8 ± 15.12(10) a</td>
<td>65.0 ± 9.30 (10) a</td>
<td>t = 6.092 <em>P</em> &lt; 0.001</td>
</tr>
<tr>
<td>3&lt;sup&gt;rd&lt;/sup&gt;/1996</td>
<td>15.7 ± 25.19(09) a</td>
<td>38.4 ± 19.66 (10) b</td>
<td>t = 2.206 <em>P</em> = 0.004</td>
</tr>
<tr>
<td>2&lt;sup&gt;nd&lt;/sup&gt;/1997</td>
<td>13.8 ± 10.48(05) a</td>
<td>64.3 ± 13.63(05) ab</td>
<td>t = 6.565 <em>P</em> &lt; 0.001</td>
</tr>
</tbody>
</table>

ANOVA: *F* = 1.897 *p* = 0.151; *F* = 4.260 *p* = 0.012

<sup>1</sup>Data transformed to arcsin (nymph mortality * 100 -1 )

Means within columns followed by the same letter do not differ significantly by the Tukey test (*P* > 0.05).

A correlation was found between the Morisita’s index and the mortality rates by predation in different generations (Pearson’s coefficient, *r* = -0.91, *n* = 4, *P* = 0.09; Fig. 5).

**Direct Observations in the Insect Communities in Pastures.** Observation of 62 nymphs showed that they predominantly (62%) disappear during the day. Based on this
preliminary observation, eight surveys of the macrofauna (insects and spiders > 2mm) in pastures were done in different hours of the day. One was done at night (7:00 to 9:00 pm) and another at the transition of sundown to the night (6:00 to 8:00 pm). All others were distributed in order to cover the whole photophase period.

The spittlebug, *D. flavopicta*, was the most abundant insect amongst species belonging to 15 taxonomic groups of herbivores and detritivores observed in the study area. That species represented 95% of observed individuals and was present in all surveys (Fig. 6). The second most common insects were Acrididae and Cicadellidae (ca. 3%). Spiders and ants were the most abundant groups of predators. Small ants such as *Pheidole*, *Solenopsis*, *Conomyrma* and *Mycocepurus* were underestimated because of their recruiting behavior (Fig. 7).

Table 2. Pearson Correlation Coefficient between the density and the mortality rate observed in each quadrat during the population peaks of the spittlebug of *D. flavopicta* during 1995/96 in the Distrito Federal.

<table>
<thead>
<tr>
<th>Generation</th>
<th>Caged</th>
<th>Uncaged</th>
</tr>
</thead>
<tbody>
<tr>
<td>1&lt;sup&gt;st&lt;/sup&gt;</td>
<td><em>r</em> = 0.334, <em>P</em> = 0.345</td>
<td><em>r</em> = 0.623, <em>P</em> = 0.054</td>
</tr>
<tr>
<td>2&lt;sup&gt;nd&lt;/sup&gt;</td>
<td><em>r</em> = 0.537, <em>P</em> = 0.110</td>
<td><em>r</em> = 0.510, <em>P</em> = 0.132</td>
</tr>
<tr>
<td>3&lt;sup&gt;rd&lt;/sup&gt;</td>
<td><em>r</em> = 0.054, <em>P</em> = 0.889</td>
<td><em>r</em> = -0.627, <em>P</em> = 0.053</td>
</tr>
</tbody>
</table>

Of the predators observed attacking or feeding upon spittlebugs, ants were the most abundant group (Table 3). Small ants (2-5 mm), like those belonging to the genera *Pheidole* and *Mycocepurus*, were observed attacking the early instars, while larger ants (10-14 mm) such as *Pachycondyla obscurocornis* Emery and *Ectatomma bruneum* Smith were observed preying upon 3<sup>rd</sup>–5<sup>th</sup> instars. In spite of an abundance of nymphs, spiders were found attacking only adults of *D. flavopicta*. This result is consistent with spiders using movement as a stimulus for recognizing prey.

**Discussion**

Our results indicate that predation is a significant factor determining *D. flavopicta*’s population density. They do so by significantly altering the population density of *D. flavopicta* immature stages, e.g. diapausing eggs and nymphs. The mortality rates of diapausing eggs were not related to population density, and did not vary significantly among years in the same pasture. However, the number of eggs presumably attacked by predators increased in relation to density. The response is consistent with a type 1 functional response, with an undefined plateau (Holling 1959 apud Hassel 1978). Our field observations showed the presence of small harvesting ants that could play an important role in the mortality of diapausing eggs. Ants of the genera *Solenopsis*, *Pheidole*, *Conomyrma* and *Cyphomyrmyx* have been reported to prey on spittlebug eggs and nymphs (Hewitt & Nilakhe 1986). The high mortality rate of *D. flavopicta* eggs, presumably by predators (ca. 60%), is similar to that observed for eggs of *Zulia entreriana* Berg. (Homoptera: Cercopidae) in pastures of *B. decumbens* in the State of Mato Grosso, Brazil (Hewitt 1986).

Mortality rates were higher in nymphal populations exposed to predators than in protected ones; this result confirms the importance of predators as a mortality factor for *D. flavopicta* nymphs. In spite of this, no preference for nymphs of *D. flavopicta* was noted among the predators. In caged cohorts, nymphs mortality rates were similar for all generations of the same year. However, in cohorts exposed to predators, mortality was significantly lower in the third and last generations of the year. We suspect that the...
predators undergo behavioral changes in response to seasonal alterations; e.g. the reduction of precipitation and day length with the progression of the wet season could be one explanation for this observation. Ants, including some species observed in our study, were reported to be more abundant in the wet season in tropical areas (Levings 1983). A shift in prey, due to the seasonal change in *D. flavopicta* nymphs abundance is another possible cause of reduction in the mortality rate (Fontes *et al.* 1995, Sujii *et al.* 1995, Begon *et al.* 1996).
Table 3. Diversity of predators observed feeding on developmental stages of the spittlebug *D. flavopicta* in pastures of *B. ruziziensis* in Brasilia, DF, Central Brazil.

<table>
<thead>
<tr>
<th>Predator</th>
<th>Prey</th>
<th>Observations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Araneae</td>
<td>Adults</td>
<td>2</td>
</tr>
<tr>
<td>One species</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Asilidae</td>
<td>Adults</td>
<td>1</td>
</tr>
<tr>
<td>One species</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Formicidae</td>
<td>Nymphs</td>
<td></td>
</tr>
<tr>
<td><em>Pachycondyla obscuricornis</em> Emery</td>
<td><em>3rd to 5th instar</em></td>
<td>3</td>
</tr>
<tr>
<td><em>Ectatomma brunneum</em> Smith</td>
<td><em>5th instar</em></td>
<td>1</td>
</tr>
<tr>
<td><em>Camponotus leydigi</em> Forel</td>
<td><em>4th instar</em></td>
<td>1</td>
</tr>
<tr>
<td><em>Labidus coecus</em> (Latreille)</td>
<td><em>3rd instar</em></td>
<td>1</td>
</tr>
<tr>
<td><em>Mycocepurus goeldii</em> Forel</td>
<td><em>1st and 2nd instar</em></td>
<td>2</td>
</tr>
<tr>
<td><em>Pheidole sp.</em></td>
<td><em>1st and 2nd instar</em></td>
<td>2</td>
</tr>
<tr>
<td><em>Pheidole gertrudae</em> Forel</td>
<td><em>1st and 2nd instar</em></td>
<td>2</td>
</tr>
<tr>
<td>Syrphidae</td>
<td>Nymphs</td>
<td>1</td>
</tr>
<tr>
<td>One species</td>
<td><em>3rd instar</em></td>
<td></td>
</tr>
</tbody>
</table>

Based on our overall data, we suggest that predation of nymphs could be acting in a density-dependent mode (as proposed by Murray 1982), and that it could influence spittlebug population levels. However, when population densities in each quadrant are analyzed independently, there is no significant correlation between nymph density and mortality rates. There was also small variation and a negative correlation between the aggregation pattern of nymphs, measured through the Morisita’s index (Sujii 1994), and the correlation between nymph density and mortality rates. There was also small variation and a negative correlation between the aggregation pattern of nymphs, measured through the Morisita’s index (Sujii 1994), and the correlation between nymph density and mortality rates. There was also small variation and a negative correlation between the aggregation pattern of nymphs, measured through the Morisita’s index (Sujii 1994), and the correlation between nymph density and mortality rates.

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